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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/462,931	01/18/2000	JUKKA HELLMAN	2328-115	5666
6449 7	7590 12/17/2002			
ROTHWELL, FIGG, ERNST & MANBECK, P.C. 1425 K STREET, N.W. SUITE 800			EXAMINER	
			COOK, LISA V	
WASHINGTON, DC 20005			ART UNIT	PAPER NUMBER
			7.1.1 0.111	
			1641	10
			DATE MAILED: 12/17/2002	Lo

Please find below and/or attached an Office communication concerning this application or proceeding.

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-4	Application No.	Applicant(s)				
	09/462,931	HELLMAN ET AL.				
Office Action Summary	Examiner	Art Unit				
	Lisa V. Cook	1641				
The MAILING DATE of this communication ap Period for Reply	pears on the cover sheet	with the correspondence address				
A SHORTENED STATUTORY PERIOD FOR REPL THE MAILING DATE OF THIS COMMUNICATION.  - Extensions of time may be available under the provisions of 37 CFR 1. after SIX (6) MONTHS from the mailing date of this communication.  - If the period for reply specified above is less than thirty (30) days, a rep - If NO period for reply is specified above, the maximum statutory period - Failure to reply within the set or extended period for reply will, by statut - Any reply received by the Office later than three months after the mailir earned patent term adjustment. See 37 CFR 1.704(b).	136(a). In no event, however, may oly within the statutory minimum of will apply and will expire SIX (6) No.e, cause the application to become	a reply be timely filed  thirty (30) days will be considered timely.  ONTHS from the mailing date of this communication.  ABANDONED (35 U.S.C. § 133).				
Status						
1) Responsive to communication(s) filed on 9/1						
,	his action is non-final.					
3) Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213.						
Disposition of Claims						
4) $\boxtimes$ Claim(s) <u>1-3,5,8,12,13 and 18-26</u> is/are pending in the application.						
4a) Of the above claim(s) <u>1-3,12,13 and 18-24</u>	is/are withdrawn from o	onsideration.				
5) Claim(s) is/are allowed.						
6)⊠ Claim(s) <u>5,8,25 and 26</u> is/are rejected.						
——7)⊠–Claim(s)- <u>5,8,25-<i>and</i>-26</u> -is/are-objected-to.						
8)⊠ Claim(s) <u>1-3,5,8,12,13 and 18-26</u> are subject	to restriction and/or elec	tion requirement.				
Application Papers						
9)⊠ The specification is objected to by the Examine	er.					
10) The drawing(s) filed on is/are: a) acce	epted or b) objected to b	y the Examiner.				
Applicant may not request that any objection to the	***					
11)☐ The proposed drawing correction filed on	_ is: a)□ approved b)□	disapproved by the Examiner.				
If approved, corrected drawings are required in re						
12) The oath or declaration is objected to by the Ex	xaminer.					
Priority under 35 U.S.C. §§ 119 and 120						
13) Acknowledgment is made of a claim for foreig	n priority under 35 U.S.0	C. § 119(a)-(d) or (f).				
a)⊠ All b)□ Some * c)□ None of:						
1. Certified copies of the priority documents have been received.						
2. Certified copies of the priority documen	ts have been received in	Application No				
3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received.						
14) Acknowledgment is made of a claim for domest	•					
a) The translation of the foreign language pro	ovisional application has	been received.				
Attachment(s)	ao priority under 00 0.0.	0. 33 120 diluiol 121.				
1) Notice of References Cited (PTO-892) 2) Notice of Draftsperson's Patent Drawing Review (PTO-948) 3) Information Disclosure Statement(s) (PTO-1449) Paper No(s)	5) Notice	w Summary (PTO-413) Paper No(s) of Informal Patent Application (PTO-152)				

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#### **DETAILED ACTION**

# Request for Continued Examination (RCE)

- 1. A request for continued examination under 37 CFR 1.114, including the fee set forth in 37 CFR 1.17(e), was filed in this application after final rejection. Since this application is eligible for continued examination under 37 CFR 1.114, and the fee set forth in 37 CFR 1.17(e) has been timely paid, the finality of the previous Office action has been withdrawn pursuant to 37 CFR 1.114. Applicant's submission filed on 13 September 2002 has been entered.
- 2. Applicants' amendment in response to the Final Office Action mailed December 13, 2001 and subsequent Advisory Action of June 7, 2002 is acknowledged (Paper #19-filed 9/13/02). In response to Amendment-F filed therein, claims 25 and 26 have been amended. Claims 5, 8, 25, and 26 are pending and currently under consideration.

#### **Drawings**

3. The drawings in this application remain objected to by the Draftsperson under 37 CFR 1.84 or 1.152 (see PTO-948). Applicant is required to submit a proposed drawing correction in reply to this Office action. However, formal correction of the noted defect can be deferred until the examiner allows the application. Applicant has deferred corrective action until allowance.

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# Information Disclosure Statement

4. The listing of references in the specification is not a proper information disclosure statement. 37 CFR 1.98(b) requires a list of all patents, publications, or other information submitted for consideration by the Office, and MPEP § 609 A(1) states, "the list may not be incorporated into the specification but must be submitted in a separate paper." Therefore, unless the examiner on form PTO-892 or applicant on form PTO-1449 has cited the references they have not been considered.

## Claim Objections

5. Claims 5, 8, 25, and 26 are objected to because of the following informalities: The dependent claims do not reference a previous claim. See MPEP 608.01(n). Appropriate correction is required.

#### Response to Argument

Applicant contends that it is customary practice to renumber the claims after allowance.

Accordingly the claimed informalities will be corrected after allowance. This argument was considered. The objection is maintained until correction after allowance.

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## Claim Rejections - 35 USC § 112

The following is a quotation of the second paragraph of 35 U.S.C. 112:

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter, which the applicant regards as his invention.

- 6. Claims 5, 8, 25, and 26 (previously 4-6 and 8-11) remain rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.
- A. Claims 25 and 26 are vague and indefinite because it is not clear what the monoclonal antibody will bind. As recited the monoclonal antibody is directed to any composition comprising the entire seq. Id. no.2 (lines 1-8 in the claims), monoclonal antibodies which bind fragments from amino acid position 7 to position 30 of seq. Id. no.2, and monoclonal antibodies which bind fragments from amino acid position 6 to position 30 of seq. Id. no.2. It is not clear if applicant intends the monoclonals of the instant invention to binding the full sequence of seg. Id. no.2 or fragments thereof?

The disclosure has support for Seq. Id. No.2, but does not clearly identify what is considered fragments that *span* position 6 to 30 with gamma-carboxylated glumatic acid positions 17, 21, and 24. In order to clearly identify the instantly claimed fragments, it is suggested that monoclonals of this type include Atcc deposit/accession numbers or seq. Id. nos. for proper identification. Please identify applicants intended meaning/define.

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## Response to arguments

Applicant contends that the claims set forth monoclonal antibodies, which bind human carboxylated osteocalcin fragments. This argument was carefully considered, but not found persuasive because the claims appear to indicate amino acids spanning position 6 to 30 of sequence identification number 2. In order to clearly identify the intended binding entity as human-carboxylated osteocalcin fragments, it is suggested that the claims be written to recite human gamma-carboxylated osteocalcin fragments of Seq. Id. No. 2. (Wherein fragments from amino acids position 7 to position 30 of seq. id. no. 2, and monoclonal antibodies, which bind from amino acid position 6 to position 30 of seq. id. no. 2 are not required). If applicant's antibodies bind at the recited amino acid positions, this should be clearly identified by the claims and reference to any human-carboxylated osteocalcin fragments, should be removed. This rejection is maintained.

#### Claim Rejections - 35 USC § 112

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

7. Claims 5, 8, 25, and 26 (previously 4-6 and 8-11) remain rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to reasonably convey to one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention.

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The written description in this case only sets forth monoclonal antibodies 2H9, 6F9, 3G8, 1C4, and 3H8 and therefore the written description is not commensurate in scope with the claims drawn to any monoclonal antibody that binds Seq. Id. No.2 (recited in independent claims 4 and 6). See pages 16-22. *Vas-Cath Inc. V. Mahurkar*, 19 USPQ2d 1111, clearly states that "applicant must convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in possession *of the invention*. The invention is, for purposes of the 'written description' inquiry, *whatever is now claimed*." (See page 1117). The specification does not "clearly allow persons of ordinary skill in the art to recognize that [he or she] invented what is claimed." (*See Vas-Cath* at page 1116). Applicant is reminded that *Vas-Cath* makes clear that the written description provision of 35 USC 112 is several from its enablement provision (see page 115).

With the exception of Mab's 2H9, 6F9, 3G8, 1C4, and 3H8, the skilled artisan cannot envision the detailed structure of the encompassed monoclonal antibodies and therefore conception is not achieved until reduction to practice has occurred, regardless of the complexity or simplicity of the method of isolation. Adequate written description requires more than a mere statement that it is part of the invention and a reference to a potential method of isolating it.

The monoclonal antibody itself is required. See *Fiers v. Revel*, 25 USPQ 2d 1601 at 1606 (CAFC 1993) and *Amgen Inc. V. Chugai Pharmaceutical Co. Lts.*, 18 USPQ2d 1016.

Furthermore, In *The Reagents of the University of California v. Eli Lilly* (43 USPQ2d 1398-1412), the court held that a generic statement which defines a genus of a compound/seq.id/etc. by only their functional activity does not provide an adequate written description of the genus.

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The court indicated that while Applicants are not required to disclose every species encompassed by a genus, the description of a genus is achieved by the recitation of a representative number of molecules, usually defined by a sequence, falling within the scope of the claimed genus.

At section B(1), the court states that "An adequate written description ...'requires a precise definition, such as by structure, formula, chemical name, or physical properties', not a mere wish or plan for obtaining the claimed chemical invention" There is insufficient description in the disclosure to support the generic claims as provided by the Interim Written Description Guidelines published in the June 15, 1998 Federal Register at Volume 63, Number 114, pages 32639-32645.

Therefore only the isolated Mab's 2H9, 6F9, 3G8, 1C4, and 3H8, but not any monoclonal that competes with the monoclonal antibodies would meet the full breadth of the claims as required by the written description provision of 35 USC 112, first paragraph.

8. Claims 5, 8, 25, and 26 (previously 4-6 and 8-11) remain rejected under 35 U.S.C. 112, first paragraph, as containing subject matter which was not described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to make and/or use the invention. The specification lacks complete deposit information for the deposit of monoclonal antibodies 2H9, 6F9, 3G8, 1C4, and 3H8.

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Because it is not clear that the properties of monoclonal antibodies 2H9, 6F9, 3G8, 1C4, and 3H8 are known and publicly available or can be reproducibly isolated from nature without undue experimentation and because the best mode disclosed by the specification requires the use of the monoclonal antibodies 2H9, 6F9, 3G8, 1C4, and 3H8, a suitable deposit for patent purposes is required. Accordingly, filing of evidence of the reproducible production monoclonal antibodies, one of ordinary skill in the art could be assured to the ability to practice the invention as claimed. Exact replication of the monoclonal antibodies is an unpredictable event.

If the deposit has been made under the provisions of the Budapest Treaty, filing of an affidavit or declaration by applicant or assignees or a statement by an attorney of record who has authority and control over the conditions of the deposit over his or her signature and registration number stating that the deposit has been accepted by an International Depository Authority under the provisions of the Budapest Treaty, that all restrictions upon public access to the deposit will be replaced if viable samples cannot be dispensed by the depository is required.

This requirement is necessary when deposits are made under the provisions of the Budapest Treaty as the Treaty leaves this specific matter to the discretion of each State. Amendment of the specification to recite the date of the deposit and the complete name and full street address of the depository is required. If the deposits have not been made under the provisions of the Budapest Treaty, then in order to certify that the deposits comply with the criteria set forth in 37 CFR §1.801-1.809, assurances regarding availability and permanency of deposits are required.

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Such assurance may be in the form of an affidavit or declaration by applicants or assignees or in the form of a statement by an attorney of record that has the authority and control over the conditions of deposit over his or her signature and registration number averring:

(a) during the pendency of this application, access to the deposits will be afforded to the Commissioner upon request;

(b) all restrictions upon the availability to the public of the deposited biological material will be irrevocably removed upon the granting of a patent on this application;

© the deposits will be maintained in a public depository for a period of at least thirty years from the date of the deposit or for the enforceable life of the patent or for a period of five years after the date of the most recent request for the furnishing of a sample of the deposited biological material, whichever is longest; and

(d) the deposits will be replaced if they should become non-viable or non-replicable.

In addition, a deposit of the biological material that is capable of self-replication either directly or indirectly must be viable at the time of the deposit and during the term of deposit.

Viability may be tested by the depository.

The test must conclude only that the deposited material is capable of reproduction. A viability statement for each deposit of a biological material not made under the Budapest Treaty must be filed in the application and must contain:

- 1)The name and address of the depository;
- 2)The name and address of the depositor;
- 3)The date of deposit;
- 4)The identity of the deposit and the accession number given by the depository;
- 5)The date of the viability test;
- 6)The procedures used to obtain a sample if the test is not done by the depository; and
- 7)A statement that the deposit is capable of reproduction.

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As a possible means for completing the record, applicant may submit a copy of the contract with the depository for deposit and maintenance of each deposit.

If the deposit was made after the effective filing date of the application for patent in the United States, a verified statement is required from a person in a position to corroborate that the cell line described in the specification as filed is the same as that deposited in the depository. Corroboration may take the form of a showing of a chain of custody from applicant to the depository coupled with corroboration that the deposit is identical to the biological material described in the specification and in the applicant's possession at the time the application was filed.

Applicant's attention is directed to In re Lundak, 773 F.2d. 1216, 227 USPQ 90 (CAFC 1985) and 37 CFR §1.801-1.809 for further information concerning deposit practice.

#### Response to Argument

Applicant argues that monoclonal antibodies have a well-documented, and highly specific, structure. And the instant antibodies have been defined by the epitope to which it binds (function). Accordingly the invention meets the requirements of written description. This argument was carefully considered but not found persuasive because even though monoclonal antibodies can be readily produced, the total characterization of a monoclonal antibody is a long and complex procedure; which varies widely with the intended use of the antibody. A general point is that if a single hybridoma has been produced and is intended for a specific function it is unlikely that the antibody produced will have all the required characteristics (Campbell, Laboratory Techniques, Vol. 13, 1984).

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Campbell teaches that it is a waste of both reagents and time to attempt full characterization of an antibody which is not obtained from a fully cloned cell line. See Chapter 10, specifically page 186. While the specification provides enough information for one of ordinary skill in the art to produce hybridoma cell lines secreting antibodies with the same or similar properties as monoclonal antibodies 2H9, 6F9, 3G8, 1C4, and 3H8, reproduction of an identical cell line and antibody is an extremely unpredictable event (see Campbell above), and because the specification lacks complete deposit information for the deposit of the hybridoma cell line(s) secreting monoclonal antibodies 2H9, 6F9, 3G8, 1C4, and 3H8, it does not appear that monoclonal antibodies 2H9, 6F9, 3G8, 1C4, and 3H8 are known and publicly available or can be reproducibly isolated from nature without undue experimentation and because certain of the claims specifically require the use of monoclonal antibodies 2H9, 6F9, 3G8, 1C4, and 3H8, a suitable deposit of the hybridoma cell lines for patent purpose is required.

Applicant has directed Examiner to the recent case of Enzo Biochem, Inc v. Gene-probe Inc. citing that the known structure and existence of a specific epitope is sufficiently descriptive. This argument was carefully considered but not found persuasive because Enzo provided reduction to practice and deposited the derived biological materials, thereby demonstrating "possession" of the invention. Applicant has not deposited the instantly claimed monoclonals, therein "possession" with respect to written description has not been meet.

Applicant contends the issue of adequate written description is dependent on the facts of each individual case and the mere citation of case law for certain broad propositions cannot be taken out of context of the specific case. Further the presently claimed invention is directed to a monoclonal antibody or recombinant fragment, which binds a specific epitope.

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This argument was carefully and fully considered but not found persuasive because the rejection was based on the facts of this individual case supported by the cited case law. The instant application being directed to monoclonal antibodies and their utility lacks written description and lack enablement because the particular inventive monoclonal antibodies have not been deposited.

Therein one of ordinary skill in the art could not be assured to the ability to practice the invention claimed. Exact replication of the monoclonal antibodies of the instant invention is an unpredictable event. The rejections are maintained.

9. For reasons aforementioned, no claims are allowed.

# Remarks

- 10. Prior art made of record and not relied upon is considered pertinent to the applicant's disclosure:
- A. Hellman et al. (Journal of Bone Mineral Research, Vol.11., No.8., 1996, pages 1165-1175) disclose nine monoclonal antibodies against osteocalcin via two-site assay procedures.

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Papers related to this application may be submitted to Group 1600 by facsimile 11. transmission. Papers should be faxed to Group 1600 via the PTO Fax Center located in Crystal Mall 1. The faxing of such papers must conform to the notice published in the Official Gazette, 1096 OG 30 (November 15, 1989). The Group 1641 Fax number is (703) 308-4242, which is able to receive transmissions 24 hours/day, 7 days/week.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Lisa V. Cook whose telephone number is (703) 305-0808. The examiner can normally be reached on Monday-Friday from 8:00 AM - 4:30 PM.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Long Le, can be reached on (703) 305-3399.

Any inquiry of a general nature or relating to the status of this application should be directed to the Group receptionist whose telephone number is (703) 308-0196.

> CHRISTOPHER L. CHIN PRIMARY EXAMINER GROUP 1800 /64/

Christych L. Chin

12/16/00

Lisa V. Cook

(703) 305-0808

12/16/02